

Figure 4 Superconducting phase displamed in Inn. The transion temperature , value defined by the once of the transion. The defined curve is a quite for the reason in the defined curve is a quite for the reason. The defined curve is a quite for the reason from electron state of the defined properties of the defined reason in the reason from electron state of the defined reason from electron state of the defined reason from electron state of the defined reason from the reason from 50 GPs. Run 4 (white the state of the defined reason of the defined reason from the reason from the reason from the defined reason from the define

of the total residual resistivity. This is because the residual resistivity includes contributions from the thin gold wire in series with the sample as well as from the finite contact resistance, which is estimated to be around 90% of the measured resistance. The main impurities contained in the starting material were O, Ta, Si and Co at concentrations of 115, 10, 83 and 8.3 p.p.m., respectively, according to the chemical analysis data. Thus we regard the drop as the onset of superconductivity of iron.

To confirm the superconductivity, we tried to detect the Meissner effect using a SQUID (superconducting quantum interference device) magnetometer at 21 GPa. A 4:1 mixture of methanol/ ethanol was used as the pressure-transmitting medium. An iron sample was obtained from the rod that provided the samples used for the resistance measurements, cut into slices 0.04 mm in thickness and 0.15 mm in diameter, and placed in the chamber of the BeCu gasket. Several turns of pick-up coal for the SQUID magnetometer were wound closely around the pressure surface of the diamond and an equal number of turns near to the diamond for background compensation. A small tin chip was put inside the compensation coal as a reference sample to check both the sign and the magnitude of the signal from iron. Figure 3 shows the superconducting transition signals from iron at 1.7 K and tin at 2.7 K, of poposite sign.

The pressure dependence of the transition temperature T, was observed in the pressure range between 15 GPa and 30 GPA (Fig. 4). Superconductivity was observed in three different runs of the resistivity measurements and the magnetization measurement. Runs 1 and 2 were obtained by quasi-hydrostatic conditions with NaCl as the pressure-medium. No pressure-medium was used for run 3 and the superconductivity transition was observed only while reducing the pressure from 90 GPa. This implies that even a very small amount of remaining ferromagnetic b.c. ciron may suppress the onset of superconductivity and that its transformation to h.c.p. iron is caused by either quasi-hydrostatic pressure or by increasing the pressure as much as possible. We conclude that the superconducting phase of iron appears after the establishment of the non-magnetic state at 15 GPa, which is close to the beginning of the box-c-to-h.c. as tructural transition of iron.

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## Direct observation of hole transfer through DNA by hopping between adenine bases and by tunnelling

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The function of DNA during oxidative stress and its suitability as a potential building block for molecular devices2-4 depend on long-distance transfer of electrons and holes through the molecule, yet many conflicting measurements of the efficiency of this process have been reported54. It is accepted that charges are transported over long distances through a multistep hopping reaction7-11; this 'G-hopping' involves positive charges moving between guanines (Gs), the DNA bases with the lowest ionization potential. But the mechanism fails to explain the persistence of efficient charge transfer when the guanine sites are distant7.12, where transfer rates do not, as expected, decrease rapidly with transfer distance. Here we show experimentally that the rate of charge transfer between two guanine bases decreases with increasing separation only if the guanines are separated by no more than three base pairs; if more bridging base pairs are present, the transfer rates exhibit only a weak distance dependence. We attribute this distinct change in the distance dependence of the rate of charge transfer through DNA to a shift from coherent superexchange charge transfer (tunnelling) at short distances to a process mediated by thermally induced hopping of charges between adenine bases (A-hopping) at long distances. Our results confirm theoretical predictions of this behaviour 15-17, emphasizing that seemingly contradictory observations of a strong as well as a weak7.12 influence of distance on DNA charge transfer are readily explained by a change in the transfer mechanism.

We have measured the efficiency of the charge transfer between Gs, separated by adenine-thymine (AT), bridges of various lengths, in double strands 1a-h (Fig. 1). Photolysis of the 4-acylated nucleotide in DNA 1 generates the sugar radical cation in 2 (Fig. 1), which injects a positive charge into G<sub>22</sub> of the complementary, radioibabled strand (2 3. Fig. 1). This guantine radical cation G<sub>22</sub> is either trapped irreversibly by water, yielding after piperidine treatment the strand cleavage product P<sub>G</sub>, or it induces an electron

(hole) transfer through DNA (3 4; Fig. 1). In order to build up a driving force for the charge transfer process, we used as electron donor a GGG triplet that is more easily oxidized than a single G<sup>II</sup>. The positive charge reaching the GGG unit was quantified by the

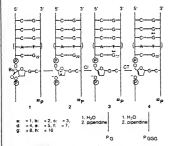


Figure 1. Charge rejection and charge transfer in DNA strainds. Busined are the method of charge protein (CI) to the GGS and orcharge transfer (CI) to the GGS sequence (S. 4), staining from DNA strainds 1a – his cross in a 4 -andystate decomparation of the Dna duce strained to the Dna duce strained as – the Dna duce strained in the double str

water trapping product P<sub>GGG</sub>, and the yield ratios P<sub>GGG</sub>/P<sub>G</sub> were determined by gel electrophoresis. We observed the product ratios shown in Fig. 2; to obtain these results, we used the assay of ref. 19 but changed the pH from 7.0 to 5.0.

The data in Fig. 2 show that the efficiency of the charge transfer, measured by the  $PGO_{2}/P_{0}$  ratio drops by a factor of 8 for each additional AT base pair in short (AT), bridges (n=1-3), But in longer sequences (n=4-7), the  $P_{GO_{2}}/P_{0}$  ratio decreases only very slightly when the number of AT base pairs increases (Fig. 2). At further elongation of the  $(AT)_{0}$  sequence, the distance influence vanishes completely. Thus, we observed no change in the  $P_{GO_{2}}/P_{0}$  ratio by increasing the number of AT base pairs from n=7 to n=16. A plot of (0)

Such a distance dependence, which demonstrates a change of the reaction mechanism, has been predicted recently 13-17. From being a coherent superexchange charge transfer (tunnelling) process at short distances, the mechanism becomes a thermally induced hopping process for long (AT), sequences, where the adenines are involved as charge carriers (A-hopping). A switch between these reaction mechanisms occurs because tunnelling rates decrease considerably as the distance increases (see the steep line at low n in Fig. 3)8.9. Therefore, in DNA strands where the guanines are separated from each other by long (AT), sequences, endothermic " transfer of the positive charge from a guanine radical cation (G \*) to an adjacent adenine becomes faster than the direct transfer of this charge to the distant guanine. The subsequent migration of the positive charge between the adenines (A-hopping) is so rapid that the length of the (AT), sequence plays only a minor role. Further experiments with DNA double strands 5 and 6 show that, in contrast to the behaviour seen with guanine radical cations, trapping of adenine radical cations by water is insignificant (Fig. 4). This difference is expected17, because the water trapping reaction of adenine radical cations proceeds through a transition state, which is much higher in energy than the transition state associated with

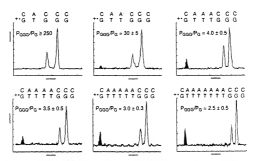


Figure 2 Ratios of the irradiation products derived from initially formed quanner ancidac abons to those derived after Circle granted across, A1 D tridges. The enduvolal curves show radioactivity interesties of indistibletion DNA strands subjected to get electronicross using operaturing polyacity and gets. They are obtained by substaction of interesties measured in control experiments (finalization of unmodified strands from acredition to irradiation permisents with the modified strands. Each. The

peaks in interest, I correspond to products P<sub>c</sub> (show no red and P<sub>coc</sub>). The peak area interest, I correspond to products P<sub>c</sub> (show no red and P<sub>coc</sub>). The peak area water area area in the peak area to produce the peak area to produce the peak area to produce the peak area cleavage of the adolesied strated Each peak area to the outcome to the peak cleavage of the adolesied strated Each peak area to the outcome to the peak area to the outcome to the peak area to the outcome to the peak area to the peak the trapping reaction of the guanine radical cation. The data of Fig 4 also confirm17 that interstrand A-hopping steps lead to a slight decrease of the hole transfer efficiency.

Our experiments show the presence of two different processes for the hole transfer between guanines in DNA: (1) a coherent superexchange reaction (single-step tunnelling), where the bridging

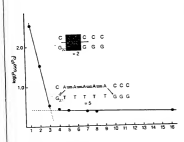


Figure 3 Plot of log(P<sub>GCO</sub>/P<sub>G</sub>) against the number of the AT base pairs. Each experiment was performed three times, and their relative errors are within 10-20% (see Fig. 2). The steep line corresponds to the coherent superexchange charge transfer. Its slope leads to ( $\beta=0.6\,\text{\AA}^{-1}$ ) of the Marcus-Levich-Jortner equation  $^{10}$  . The flat line is drawn in order to make clear the weak distance dependence. The product ratios P<sub>009</sub>/P<sub>Q</sub> are proportional to the charge transfer rates. The arrows in the depicted DNA strands indicate the superexchange charge transfer between G<sub>22</sub> and the GGG sequence for short distances

( = 2), or the A-hopping mechanism for long distances ( = 5), where---in addition--adenines act as charge carners. For clarity, only the double strands with = 2 and = 5 are shown. The nucleotides in red indicate all charge carriers.

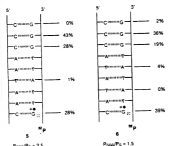


Figure 4 Relative yields of water trapping products during hole transfer through DNA sequences 5 and 6. The experiments were performed three times (strand 5) or twice (strand 6). The relative errors of yields of products P<sub>GGG</sub> and P<sub>G</sub> are 15%. The yields obtained by trapping the A sites have absolute errors of 4%. Trapping products of thymine bases could not be detected. The nucleotides in red indicate the beginning and the end of the charge transfer process.

adenines are indirectly affecting the transfer mechanism by mediating the electronic coupling between the guanines, and (2) a thermally induced hopping process (A-hopping mechanism), where the lifetime of the guanine radical cation is long enough to oxidize the intervening adenine bases and directly involve them in charge transport. The efficiency of the tunnelling reaction decreases rapidly with the number of the intervening AT base pairs, whereas the A-hopping process is only slightly influenced by the number of the AT base pairs. Thus, the debated contradiction between very strong to and very weak 7.12 influence of distance on the hole transfer through DNA can be explained by a change in the reaction mechanism.

These results indicate that A-hopping provides a mechanism for ameliorating the harmfulness of damage to DNA under conditions of oxidative stress, by providing a means for charge transport over long sequences of AT pairs into regions of low ionization potentials<sup>1,21</sup>. In addition, the ability to rationally change the behaviour of a system from strongly distance-dependent charge transfer to weakly distance-dependent charge transfer may open new opportunities in nanoelectronics. DNA can offer this ability because the switch to the activated hopping mechanism depends on the difference of the oxidation potentials of the DNA bases involved-and this difference could be easily changed by using suitable substituents.

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Perc/Pc = 2.5